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DOI: <https://doi.org/10.1093/nar/gkz1151>

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ZORA URL: <https://doi.org/10.5167/uzh-182692>

Journal Article

Published Version



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Originally published at:

Voith von Voithenberg, Lena; Fomitcheva Khartchenko, Anna; Huber, Deborah; Schraml, Peter; Kaigala, Govind V (2020). Spatially multiplexed RNA in situ hybridization to reveal tumor heterogeneity. *Nucleic Acids Research*, 48(3):e17-e17.

DOI: <https://doi.org/10.1093/nar/gkz1151>

Spatially multiplexed RNA *in situ* hybridization to reveal tumor heterogeneity

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Received October 09, 2019; Revised November 20, 2019; Editorial Decision November 21, 2019; Accepted December 03, 2019

ABSTRACT

Multiplexed RNA *in situ* hybridization for the analysis of gene expression patterns plays an important role in investigating development and disease. Here, we present a method for multiplexed RNA-ISH to detect spatial tumor heterogeneity in tissue sections. We made use of a microfluidic chip to deliver ISH-probes locally to regions of a few hundred micrometers over time periods of tens of minutes. This spatial multiplexing method can be combined with ISH-approaches based on signal amplification, with bright field detection and with the commonly used format of formalin-fixed paraffin-embedded tissue sections. By using this method, we analyzed the expression of *HER2* with internal positive and negative controls (*ActB*, *dapB*) as well as predictive biomarker panels (*ER*, *PgR*, *HER2*) in a spatially multiplexed manner on single mammary carcinoma sections. We further demonstrated the applicability of the technique for subtype differentiation in breast cancer. Local analysis of *HER2* revealed medium to high spatial heterogeneity of gene expression (Cohen effect size $r = 0.4$) in equivocally tested tumor tissues. Thereby, we exemplify the importance of using such a complementary approach for the analysis of spatial heterogeneity, in particular for equivocally tested tumor samples. As the method is compatible with a range of ISH approaches and tissue samples, it has the potential to find broad applicability in the context of molecular analysis of human diseases.

INTRODUCTION

Personalized medicine relies on the molecular analysis of individual tumors. To understand differences in progression and survival probabilities between patients affected by cancer, it is necessary to differentiate tumors into subtypes. For example, the current molecular classification schemes

for breast cancer describe five molecular subtypes: luminal A, luminal B, human epidermal growth factor receptor 2 (HER2)-positive, triple-negative, and normal-like (1). For predictive and prognostic reasons, breast tumors are classified into these subtypes according to their molecular characteristics. Standard clinical methods for subtype differentiation include immunohistochemistry (IHC) to detect the biomarker expression status at the protein level and fluorescence *in situ* hybridization (FISH) to analyze gene amplification. While most of the tumors can be classified by using these methods, equivocal cases do exist. For HER2 status determination, for example, 32% of tumors are described as equivocal when using IHC and 5% when using FISH (2). Repeated testing using both methods can reduce the total number of equivocal cases to 5%. Ohlschlegel *et al.* showed that equivocal testing is largely influenced by the molecular heterogeneity of the tumor tissue (3). While genomic heterogeneity was described for 14.7% of the cases in a general cohort, the number increased to 41% for equivocal cases. The analysis of molecular heterogeneity is therefore highly valuable for the subtype differentiation of breast tumors and thus also for prognostic and predictive purposes.

Cellular heterogeneity is a driving factor in development and disease. It arises from modulations at the gene, transcript, and protein level and can be detected molecularly and phenotypically. Patient-by-patient or intertumor molecular heterogeneity is commonly analyzed in the clinical setting (subtype differentiation) and can be distinguished and addressed by using existing methods recommended in clinical guidelines (4). However, the impact of the prognostic and predictive power of spatial intratumor heterogeneity is less known. Retaining spatial information is highly important to link the molecular state to function, development, and disease. This is of special significance in tumors, as ~60–70% of all somatic mutations are not detectable in all tumor areas (5) and cell variants with an abundance of as little as 0.1% of the total cell population can affect the treatment response (6). Recently, the coexistence of multiple breast cancer subtypes and their plastic conver-

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sion within a single tumor has been proposed (7) and several clinical trials are currently analyzing the impact of intratumor heterogeneity spatially and temporally (8–11). Interestingly, intermediate levels of genetic intratumor heterogeneity have been linked to a poor prognosis (12,13), while, for high levels of heterogeneity, the prognosis improved in certain cases (12). To personalize tumor treatment based on heterogeneity profiles, it is therefore important to capture the molecular tumor profile to the full extent in the spatial and temporal dimension. *In situ* techniques intrinsically provide spatial information and *in situ* protein detection has been extended to multiplexed approaches, thus allowing the spatial detection of multiple proteins of interest. In tumor analysis, multiplexing is commonly used for the detection of protein biomarkers, e.g. using the Opal multiplex assay (14), and tumor associated immune cells (15–17). Through using automated microfluidic systems, the duration of fourplex experiments has been reduced from two days to few hours (14,18). Quantitation of the signal is facilitated by immunofluorescence approaches and spectral unmixing or sequential detection (19–21) and can be performed using automated quantitative analysis (AQUA) (22). Nonetheless, when using current standard immunohistochemistry approaches (23), the quantification of protein expression levels is difficult (24). Therefore, the analysis and quantification of transcripts is a promising alternative, as it provides information about the genetic content as well as gene expression levels (25). Recent novel diagnostic developments for breast cancer thus place a focus on the detection of gene expression on the RNA level, e.g. 21-gene Oncotype DX Recurrence Score (26), 70-gene MammaPrint Assay (27) and a multitude of other signatures (28,29), as mRNA is readily quantifiable in quantitative real-time polymerase chain reactions (qRT-PCR) or microarray-based approaches. Clinical guidelines approve of the use of gene expression profiles for additional prognostic and predictive purposes under certain conditions (29,30) and ongoing studies of tumors in the central nervous system show great promise in using RNA sequencing as a predictive measure (31). Translating such methods for quantifying gene expression from the research to the clinical setting therefore plays an important role in developing novel diagnostic approaches for addressing intratumor heterogeneity.

In research settings, several techniques for the detection and quantification of transcripts exist. While many of the early quantitative methods required the extraction of RNA from the sample (such as Northern blotting, qRT-PCR, digital PCR, microarrays or high-throughput sequencing), new developments allow for *in situ* detection and quantification of transcripts. These methods can be divided into four categories: (I) FISH (32–36), (II) FISH with barcoding (37–40), (III) *in situ* sequencing (41–44) and (IV) patterned barcoded microarrays (45,46). These methods have in common that they provide information about transcript localization in the sample or even on a subcellular level. They thus enable the analysis of subpopulations of cells, whose signals would be averaged in extraction-based approaches. A methodology combining the advantages of both, localization-retaining methods and high-throughput sequencing, uses laser capture microdissection (47–49). While spatial transcriptomic methods, such as *in*

situ sequencing or combined laser capture microdissection and RNA sequencing, provide quantitative spatial information about a large number of transcripts (up to 6000), they require the use of specialized instruments and techniques, which are currently not used in the clinical environment. As DNA-FISH is routinely performed in the clinical setting, the equipment required for *in situ* hybridization experiments is available and can be readily used for transcript analysis. Wang *et al.* described the use of RNA *in situ* hybridization as a quantitative companion diagnostic application for HER2 testing, with special importance for cases with intratumoral heterogeneity or equivocal FISH results (2). Previous studies have shown the applicability of RNA-ISH for formalin-fixed paraffin-embedded (FFPE) tissue sections (50–52). Compared to other spatial transcriptomic techniques, it thus poses an important alternative for clinical applications because the majority of tumor tissues stored in biobanks are FFPE tissues. Furthermore, nearly all mRNA molecules can be detected using single molecule FISH, while single-cell mRNA sequencing captures 5–40% and microarray or *in situ* barcode sequencing only around 5% of mRNA molecules (34). This makes RNA-ISH a wide-spread technique for quantitative or semi-quantitative analyses of transcripts, although limited in the number of species that can be detected simultaneously. The detection of single RNA transcripts and their quantification can either be performed directly, by using hybridization probes containing fluorophores and highly sensitive detection optics (53), or by amplification of the reaction using branched FISH (54–56), enzymatic processes (57) and antibodies (58). The use of amplification-based systems obviates the need for expensive optical setups and detection can even be performed using brightfield microscopy. Its application is currently limited, however, to one or two detection channels depending on the enzyme-substrate pairs used. We therefore use a microfluidic system for the local delivery of probes, which allows for spatial multiplexing. We make use of a variant of a microfluidic system called the vertical microfluidic probe (MFP) (59,60) in this work, which enables the localized delivery of reagents on the nanoliter scale. To avoid confusion with the usage of FISH probes, we term the MFP as microfluidic chip in this manuscript. A basic implementation of the chip consists of two channels with a size in the range of 50–100 μm , whose openings are positioned in parallel to and tens of micrometers above a surface of interest (59,60). Simultaneous injection and aspiration of liquid (at certain ratios of flow rates) from the channels confines the injected reagent to a small region between the chip and a surface immersed in aqueous solution (hydrodynamic flow confinement). Owing to its compatibility with biological substrates in a liquid environment, variants of the microfluidic chip (60) have found use in life science and biomedical applications (24,61,62).

Here, we established a microfluidic method for the detection of multiple transcripts in FFPE tissues through using spatial multiplexing in branched probe-based or enzymatic amplification RNA-ISH. Even though current clinical approaches are based on the semi-quantitative analysis of proteins and DNA, the possibility of quantification on the transcript level will likely provide advantages in the description of the spatial heterogeneity of cancers,

where gene amplification, RNA and protein levels correlate (2,63). By using this multiplexed RNA-ISH, transcripts of interest along with tissue internal controls, complex cancer biomarker panels and spatial tumor heterogeneity can be evaluated on a single tissue section.

MATERIALS AND METHODS

FFPE cell blocks and tissues

FFPE cell blocks of the cell lines MCF7, BT474 and SKBR3 (AMS Biotechnology, Abingdon, UK) with HER2 expression levels of 1+, 2+ and 3+, respectively, were used as references. Breast carcinoma tissue sections were either obtained from BioChain (T2235086) or were kindly provided and anonymized by the tissue biobank of the Department of Pathology and Molecular Pathology of the Universitätsspital Zürich.

Hematoxylin and eosin staining of FFPE sections

Tissue sections were stained by incubation in hematoxylin for 5:30 min. A subsequent washing process in tap water for 5 min, incubation in 1% HCl for 10–20 s, and another wash in tap water for 1 min were used for colorimetric changes and removal of excess hematoxylin. Eosin was applied for 1 min, followed by a wash in tap water for 10 s and posterior dehydration of the tissue section using 95% and 100% ethanol for 1 min each.

Microfluidic system for probe delivery

The microfluidic chip, consisting of one layer of silicon and one of glass and containing four channels (channel size: 200, 100, 100, 200 μm) was fabricated in clean-room facilities in-house (60) and is reusable under stringent cleaning conditions. It was positioned on the microscopic system on a stage adjustable along the z-axis, using a connector system consisting of a screw, a gasket ring and a circular connector (Dolomite Microfluidics) (Supplementary Figure S1A–C). Tubings emerging from four syringes serving as reservoirs and a flow system were attached to the microfluidic chip via the connector. Liquid flows were generated using NeMESYS low pressure module syringe pumps (Cetoni, Korbussen, Germany) controlled by the software QMix Elements (Supplementary Figure S1D). The microfluidic chip was mounted on a Nikon Ti Eclipse microscope body placed in an incubating chamber (Life Imaging Services) and equipped with a Lumencor Sola 6-LCR-SB illumination (Lumencor, Beaverton, USA) and a Hamamatsu Orca Flash 4.0 camera (Hamamatsu, Nakaku, Japan). The position of the sample was adjusted in the x- and y-direction using high-precision stages (Phytron, Gröbenzell, Germany) controlled by the software WinCommander5.

RNA *in situ* hybridization

RNA-ISH experiments were performed following the instructions of the manufacturer (ViewRNA, Thermo-Fisher Scientific). Solutions for hybridization and washing were provided as part of the *in situ* hybridization kit. ISH probes targeted against *dapB* (*dapB* ViewRNA Probe Type 6,

Bacillus subtilis), *HER2* (HER2 ViewRNA Probe Type 1), *PgR* (PGR ViewRNA Probe Type 1 and Type 6), *ER α* (ESR ViewRNA Probe Type 1 and Type 6) and *Actin β* (ActB ViewRNA Probe Type 1) RNA were used.

The FFPE tumor sections were incubated at 60°C for 1 h prior to the removal of paraffin using a 10 min incubation in xylene, two 5 min incubations in ethanol and washing steps in 70% ethanol and H₂O. The pretreatment of the sections was a 20 min incubation of the slides in a 1 \times preincubation solution at 95°C. While incubating the sections in PBS, a hydrophobic pen was applied (ImmEdge Pen, Vector Laboratories) to limit the spread of the immersion liquid to a defined area around the tumor section. The sections were then digested using protease at 40°C for 5 min (cell pellet sections) or 10 min (mammary carcinoma sections). After washing in PBS, the sections were fixed using 10% neutral buffered formalin solution for 5 min. Sections were washed in PBS before the primary probe was delivered either through using the microfluidic chip (see *Delivery of primary ISH-probes using the microfluidic chip*) or by incubation of the whole slide in probe solution at 40°C. Varying hybridization times of the primary probe between 10 min and 3 h were used as indicated. After washing the sections in wash buffer three times, they were incubated with preamplifier solution (for dual color experiments: preamplifier mixture) at 40°C for 25 min. Following three washes in wash buffer at room temperature, the sections were incubated in amplifier solution (for dual color experiments: amplifier mixture) at 40°C for 15 min. Residual probes were removed by washing the sections three times using wash buffer. When dual-color experiments were performed, the sections were incubated with a label-6 alkaline phosphatase (AP) probe at 40°C for 15 min, washed three times, and incubated at room temperature for 30 min using a blue reagent as enzymatic substrate. After three washes using wash buffer, residual alkaline phosphatase was deactivated using AP stop solution at room temperature for 30 min, and the sections were washed using PBS and wash buffer. The sections were then incubated with a label-1 alkaline phosphatase probe at 40°C for 15 min, residual probes were removed using wash buffer, and the reaction was enhanced by an incubation in AP enhancer solution at room temperature for 5 min. The sections were incubated with Fast Red substrate at 40°C for 30 min, before being stained using hematoxylin and 3 $\mu\text{g}/\text{ml}$ 4'-6-diamidino-2-phenylindole (DAPI). Sections were mounted using aqueous ImmunoHistoMount (Sigma-Aldrich) and imaged using brightfield and fluorescence detection (see *Signal Visualization*). Using CellProfiler (64), nuclei and cytoplasm of cells were detected and the ISH signal intensity was determined and allocated to each of the cells.

Delivery of the primary ISH-probes using the microfluidic chip

We performed hematoxylin and eosin staining on sections consecutive to the ones used for local RNA-ISH. Regions for delivery of the ISH probes were defined by microscopic inspection of the hematoxylin and eosin stained tissue sections. Primary probes were delivered using a microfluidic chip positioned ~ 50 μm above the tissue section immersed

in PBS. The inner channels of the chip with channel widths of 100 μm were used for the delivery of the primary probe, while the outer channels (channel width 200 μm) were used for the visualization of the hydrodynamic flow confinement using a red food dye (Trawosa). The flow rates of the inner and outer injections were 1 $\mu\text{l}/\text{min}$ and 0.5 $\mu\text{l}/\text{min}$ and the ones of the inner and outer aspirations were 1 $\mu\text{l}/\text{min}$ and 5 $\mu\text{l}/\text{min}$, respectively. When performing the experiments using the microfluidic chip, primary probe incubation times ranged from 10 min to 1 h. The positions of local delivery of the probes were recorded using the software of the x-y-z stages (WinCommander5).

Whenever multiplexed RNA-ISH experiments of a single detection color and involving the use of primary probes of type 1 and type 6 were performed using the microfluidic chip, preamplifier and amplifier mixtures were used and the label probes of type 1 and type 6 were combined in a single hybridization reaction.

Global and microfluidic chip-based immunohistochemistry

ER, PgR and HER2 detection and quantification through immunohistochemistry was performed following the guidelines of the manufacturer (Herceptest, Dako Denmark A/S) and primary antibodies were delivered either globally or locally using the microfluidic chip as described (24,59). Briefly, after deparaffinization, epitope retrieval was performed at 95–99°C for 40 min (+20 min cool-down). After washing in wash buffer, the sections were incubated in a peroxidase-blocking solution for 5 min. For global immunohistochemistry, the sections were incubated in 2 $\mu\text{g}/\text{ml}$ anti-HER2 antibody (ErbB2 Monoclonal Mouse Antibody, Clone e2-4001, ThermoFisher) for 1 h, in a 1:40 dilution of anti-ER antibody (Monoclonal Rabbit Anti-Human Estrogen Receptor α , Clone EP1, Dako) for 20 min, or in a 1:50 dilution of anti-PgR antibody (Monoclonal Mouse Anti-Human Progesterone Receptor, Clone PgR 1294, Dako) for 20 min. For local immunohistochemistry, 20 $\mu\text{g}/\text{ml}$ anti-HER2 antibody (Polyclonal Rabbit Anti-Human c-erbB-2, Dako) was delivered to the tissue sections using the microfluidic chip for incubation times ranging from 28 to 600 s. The sections were then washed in wash buffer and incubated with a visualization reagent containing secondary antibodies against mouse and rabbit (EnVision Dual Link System-HRP, Dako) for 30 min. After washing the sections, the signal was visualized using 3,3'-diaminobenzidine for 10 min. A hematoxylin counterstain was performed to visualize the cells.

Signal visualization and analysis

Brightfield and fluorescence imaging was performed on a Nikon Eclipse 90i microscope using 4 \times 0.2 NA, 10 \times 0.45 NA, 20 \times 0.75 NA, and 40 \times 0.6 NA objectives. Brightfield signals were detected on a DXM 1200C camera.

For fluorescence imaging, the sample was excited using a Lumencor Sola SE 5-LCR-SB with the excitation filters 531/40 nm or 630/38 nm for visualization of the Fast Red or the Fast Blue signal, respectively. Fluorescence was collected by 4 \times 0.2 NA, 10 \times 0.45 NA, 20 \times 0.75 NA and

40 \times 0.6 NA objectives and separated according to wavelength by a 562 nm dichroic combined with a 593/40 nm emission filter or a 757 nm dichroic and a 775/59 nm emission filter. Stained nuclei were visualized using a DAPI filter set (377/50, 409, 447/60 nm). The signal was detected on a Digital Sight DS-1QM camera. Signal intensities were extracted from the images as integrated intensities of single cells using CellProfiler (64).

Heterogeneity analysis

The extracted signal intensities were compared between different regions of the tumor tissue section and a Mann-Whitney U -test was used to determine differences in the expression levels between regions. The test statistic U was calculated as

$$U = n_1 n_2 + \frac{n_1(n_1 + 1)}{2} - R_1,$$

where R_1 is the larger of the two rank sums, n_1 the number of samples in the group with the larger rank sum and n_2 the number of samples in the group with the smaller rank sum. The significance z was standardized using

$$z = \frac{U - \mu_U}{\sigma_U}$$

with the mean μ and the standard error σ of the distribution U . The Cohen effect size r , describing the importance of the significance in the difference of the means, was determined as

$$r = \left| \frac{z}{\sqrt{n}} \right|,$$

where n is the sum of the number of samples in both groups. A Cohen effect size $r = 0.1$ typically corresponds to a small effect, $r = 0.3$ to a moderate effect, and $r = 0.5$ to a strong effect (65). For visualization, the effect size was displayed as a color-coded matrix.

RESULTS

Localized RNA-ISH by microfluidic implementation

The molecular heterogeneity of tumor tissues is reflected in spatial differences in gene expression. To investigate these differences at the transcript level, we developed a microfluidic implementation of the RNA-ISH assay (Figure 1A, Supplementary Figure S1). It makes it possible to increase the number of transcripts that can be investigated on a single tissue section by using amplification-based transcript detection. Theoretically, up to 3000 hybridizations could be performed on a tissue area of 10 mm \times 10 mm. However, the number of reactions that one can practically perform depends on multiple factors, ranging from the morphology of the tissue to availability of probes and design of the microfluidic chips.

The initial pretreatment of the tumor tissue section, including paraffin removal, heat-induced pretreatment and protease digestion, was performed on the whole tissue slide. Precise control of the pretreatment steps, especially the protease digestion times, specific to the tissue of origin was re-

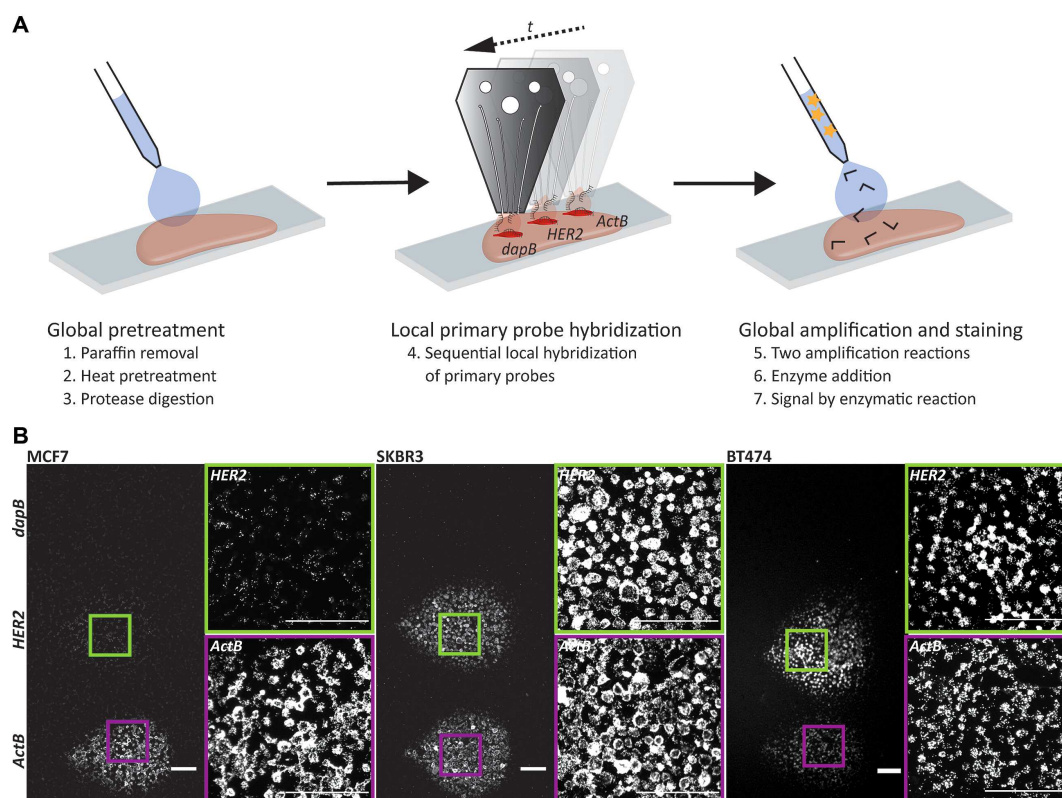


Figure 1. Local RNA *in situ* hybridization. (A) Schematic overview of a microfluidic implementation of chromogenic RNA *in situ* hybridization. The initial preparation of the tissue section including paraffin removal (1), heat (2) and protease (3) pretreatment, as well as fixation was performed globally. The primary probe was then hybridized locally to the RNA of interest (4). In this case the detection of *HER2* is shown with internal negative (*dapB*) and positive (*ActB*) controls. Finally, the amplification of the signal using preamplifier and amplifier strands (5), the binding of enzyme (6), and the enzymatic reactions (7) were performed globally on the whole slide. (B) Fluorescence images of RNA-ISH experiments detecting the gene expression of *HER2* in cell pellet sections of the breast cancer cell lines MCF7 (left), SKBR3 (center) and BT474 (right). Probes directed against bacterial *dapB* and *ActB* were used as negative and positive controls, respectively. Scale bar: 100 μ m.

quired (Supplementary Note SN1, Figure S2). For the subsequent delivery of the primary probe, we used an open space microfluidic chip (59,60,66). Through the simultaneous injection and aspiration of the primary probe solution, it was delivered to a spatially defined region of ~ 50 – 100 cell sections (~ 150 μ m in diameter). Probes for multiple transcripts of interest can thus be used in neighboring regions within the same tumor area. Additionally, through using hierarchical flow confinements (66), inner and outer regions of the confinement can be targeted using different probes and thus patterns of local delivery can be generated. The RNA-ISH signal increased for primary probe delivery times from 10 to 60 minutes, even under conditions with a constant flow (Supplementary Figure S3). Further increasing the hybridization time of the primary probe beyond 60 min, however, did not increase the signal intensity. Therefore, an incubation time of 60 min was chosen, when using the microfluidic chip. Constant delivery of probes under flow leads to the convection-enhanced reduction in hybridization time, which facilitates the sequential delivery of multiple probes without affecting the time required for the complete workflow (11–14 h depending on the spatial and spectral multiplexing conditions). Subsequent signal amplification and visualization was performed on the whole tumor tissue section, which enables direct comparisons of the

signals within each tumor tissue section without having to take additional properties of the fluorophores and detection optics into consideration.

The use of the microfluidic chip system thus allows for the spatial multiplexing of a number of different RNA targets. The global delivery of oligonucleotides for amplification of the signal and enzymatic substrates permits different targets to be simultaneously amplified and visualized in a single reaction, thereby facilitating quantitative comparisons that are independent of the amplification process.

Biomarker analysis with intrinsic positive and negative controls

The use of the microfluidic chip for spatially defined RNA-ISH experiments enables the sequential or even simultaneous deposition of primary probes targeted at several RNA sequences of interest on a single tissue section. We leveraged this ability to introduce probes for the biomarker of interest, *HER2*, as well as 4-hydroxy-tetrahydronicotinate reductase (*dapB*) from *Bacillus subtilis* and *Actin* β (*ActB*) as negative and positive controls on the same slide. The positive control serves as an indicator for RNA integrity, while the negative control, which is not expressed in human tissue, provides information about background enzymatic

reactions, unspecific binding of amplification probes, and staining. The absence of *dapB* signal in the area of local delivery and the absence of ISH signal in cells surrounding the spatially defined regions serve as negative controls. Figure 1B shows FFPE cell pellet sections of the cell lines MCF7, SKBR3 and BT474 with RNA-ISH signals of *dapB*, *HER2* and *ActB* in spatially defined regions of the section (see also Supplementary Figures S4). The relative quantification of the signal intensity was based on the mean integrated intensity of the FISH signal per cell section (Supplementary Figure S5). The relative comparison of expression levels within a tissue section is independent of the amplification, enzymatic visualization reaction, and imaging conditions, as these are equal to all transcripts of interest in the same section. The expression levels based on ISH signal intensities were therefore directly compared within a single tissue section. The signal of the positive control *ActB* was detected for all three cell lines with a mean integrated intensity per cell between 40 and 100, while the *dapB* signal was low in all cases (mean integrated intensity of 3–10) (Figures 1B, 2). The signal intensity of *HER2* showed a strong variation between the three cell lines with a high mean integrated intensity per cell of 438 for SKBR3 and 208 for BT474 and a weak signal (int = 7) for MCF7. This is consistent with the expression levels detected for these cell lines when using immunohistochemistry, where SKBR3 and BT474 are described as HER2 3+ and MCF7 as HER2 0–1+ (Supplementary Figure S6) (67). The method thus enables the estimation of HER2 overexpression in known breast cancer cell lines. Therefore, we applied the same method to investigate the *HER2* expression in breast carcinoma sections and found *HER2* to be strongly expressed when compared to the positive control *ActB* in the tumor region analyzed (Figure 2A). The analyzed tumor is thus HER2 positive.

Using this approach, the comparison to a house-keeping gene, such as *ActB*, can be used to quantify differences in the expression level of the gene of interest (68). We calculated the ratios of *HER2* and *ActB* expression levels based on the mean integrated intensities per cell section. For MCF7 cells, a ratio of 0.07 was calculated. BT474 and SKBR3 expressed *HER2* and *ActB* in ratios of 5.65 and 6.64, respectively. The investigated mammary carcinoma section showed an expression ratio of 23. However, as shown by Aerts *et al.* (69), we observed that the expression level of *ActB* at the transcriptomic level, based on the integrated intensity of the ISH signal, differs between the breast cancer cell lines MCF7, SKBR3 and BT474 (Figure 2B, Supplementary Note SN2). A direct comparison between the different cell lines therefore has to be considered carefully when basing the quantification on the house-keeping gene *ActB*. While different cell cycle states within a certain cell line may indeed be compared, additional house-keeping genes should be considered for the comparison between cell lines or tumor tissues.

We demonstrated here that transcripts of interest can be analyzed in a multiplexed fashion using the microfluidic chip-based delivery of ISH probes. Nonetheless, house-keeping genes used for the relative quantification of transcript levels must be evaluated carefully. Integrating negative and positive controls into the experiments provides measures for unspecific signal and ensures that pre-

analytical and analytical conditions (e.g. fixation, pretreatment of the tissue, assay parameters) do not impact RNA integrity. Our approach thus provides a robust basis for a controlled gene expression analysis.

Multiplexed RNA-ISH analysis of a predictive breast cancer biomarker panel

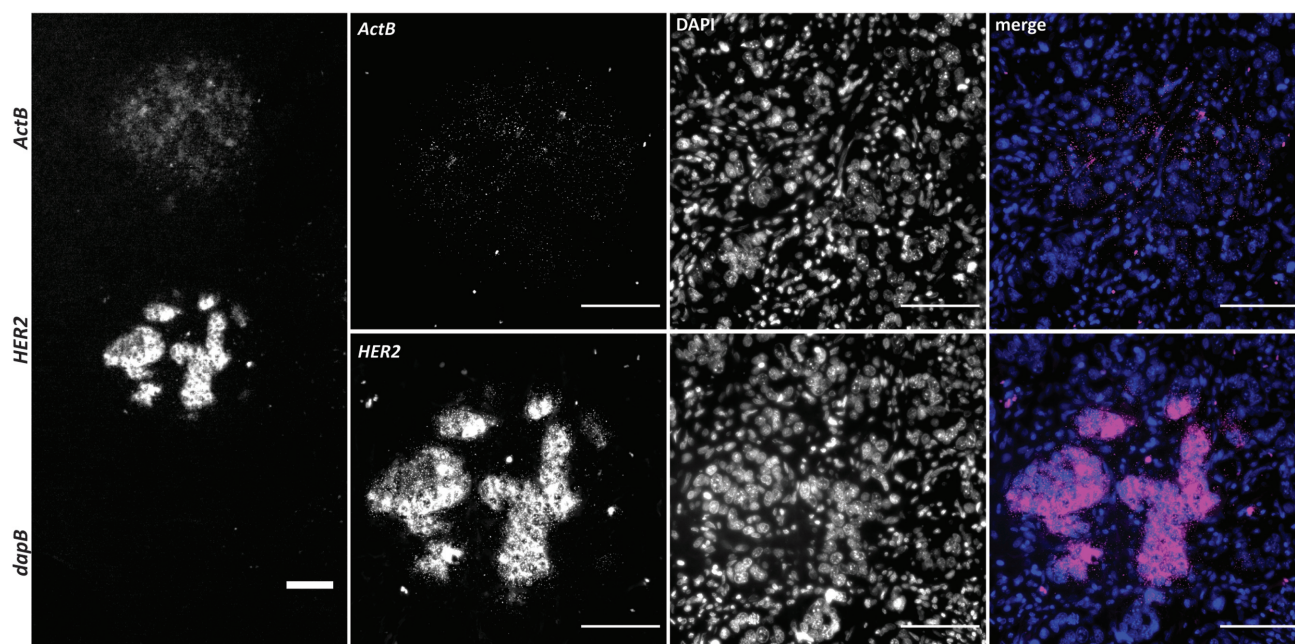
Spatially multiplexed RNA-ISH allows the analysis of biomarker panels using single color detection. We focused the analysis on the predictive breast cancer biomarker panel estrogen receptor (*ER*), progesterone receptor (*PgR*), and *HER2*. By using microfluidic chip-based local delivery, we determined their expression levels in cell pellet sections of breast cancer cell lines and mammary carcinoma sections (Figure 3, Supplementary Figures S7). The imaging conditions were kept constant for the analysis of the different cell lines and the carcinoma tissue in Figure 3. We observed an elevated expression of *ER* in MCF7 (int = 29) and *PgR* in MCF7 and BT474 cells (int = 16 and int = 535, respectively). *HER2* was strongly overexpressed in SKBR3 (int = 1057) and BT474 (int = 836) and mildly expressed in MCF7 (int = 20) (Supplementary Figure S7). The mammary carcinoma section showed *HER2* overexpression, while *ER* and *PgR* signals were not elevated (Figure 3). These results correspond to the ones of the molecular analysis of the cell lines by immunohistochemistry. In MCF7, the Allred scores for *ER* and *PgR* are 6, whereas *HER2* expression is quantified as 0–1+ (67). SKBR3, on the other hand, shows a high expression of *HER2* (3+) and no overexpression of *ER* and *PgR* (Allred score = 0) and BT474 highly expresses *HER2* (3+) and *PgR* (Allred score = 8) (67).

These results show that the multiplexed analysis of multiple transcripts, e.g. a biomarker panel, is possible through using single-color RNA-ISH in fluorescence or brightfield detection mode. We believe that this facilitates the quantification of multiple transcripts at the same time and allows for the detection of a multitude of transcripts requiring only a single tissue section or small amounts of valuable tissue.

We further increased the multiplexing modalities by combining spatial multiplexing with dual color detection. By using only two spatial positions, we could thus visualize gene expression patterns of the biomarker panel *ER/PgR/HER2* and the positive control *ActB* as well as negative controls in forms of the surrounding tissue (Supplementary Figure S8). *ActB* and *HER2* were visualized by using Fast Red, and *ER* and *PgR* using Fast Blue. The *ActB* signal and the area surrounding the local delivery of probes show the integrity of the RNA in the sections as well as a negligible signal owing to unspecific reactions. As expected, MCF7 showed an overexpression of *ER* and *PgR*, while the *HER2* signal was significantly lower than for the other cell lines. On the other hand, SKBR3 and BT474 as well as the carcinoma sections, showed a strong *HER2* signal. *PgR* was dominantly overexpressed in BT474 cell pellet sections. This agrees with the subtype differentiation of these cell lines into luminal A, luminal B, and *HER2*-overexpressing with an Allred score of 8 for *PgR* in BT474 and one of 6 in MCF7 cells.

Although the sensitivity of detection is higher when using fluorescence excitation and detection, the spatial multiplexing approach allows the detection of multiple different

A



B

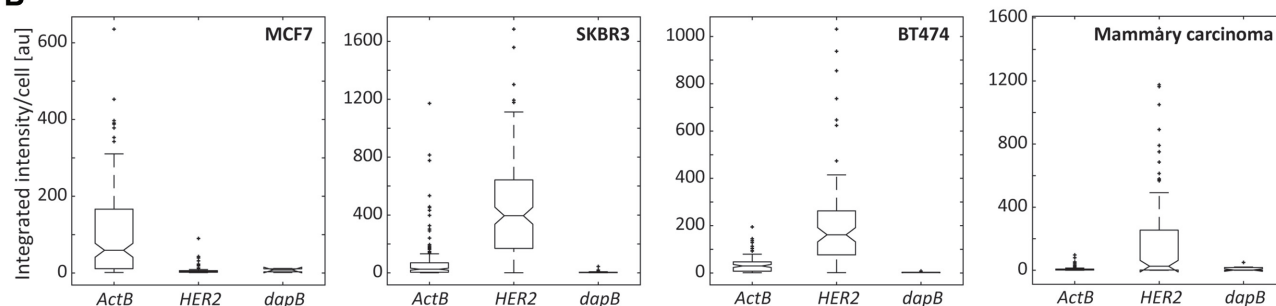


Figure 2. Detection of *HER2* expression status by RNA *in situ* hybridization using internal positive and negative controls. (A) Fluorescence images of the signals detected for *HER2*, *dapB*, and *ActB* in a mammary carcinoma section. Scale bar: 100 μm . (B) Quantitative analysis of the fluorescence intensity of FISH signals detected for *HER2*, *dapB*, and *ActB* integrated per sectioned cell for FFPE sections of the breast cancer cell lines MCF7, SKBR3, and BT474 (see Figure 1) as well as the mammary carcinoma from (A). 100 ms excitation.

types of transcripts when using brightfield microscopy on a single tissue section (Supplementary Figures S4, S9). By using micrometer sized areas of hybridization, thousands of transcripts could theoretically be detected on a single tissue section through spatial multiplexing. By additionally employing dual-color detection schemes, the number of different types of detected transcripts can be doubled.

Spatially multiplexed RNA-ISH for detecting molecular tumor heterogeneity

Having shown that multiplexed RNA-ISH detection methods allow for the spatial analysis of several biomarkers simultaneously, including internal positive and negative controls (Figure 2) or panels of biomarkers (Figure 3), we applied the technique to investigate spatial heterogeneity in the expression levels of the detected transcripts. By using globally or locally applied RNA-ISH, we analyzed the spatial heterogeneity of *HER2* expression in mammary carcinoma sections (Figure 4, Supplementary Figure S10). Each

tumor region (distance between regions 5 mm to 1 cm, separated by fibrous tissue) was represented by multiple positions within the region (areas with a distance of a few 100 μm). Differences within each of the regions (intraregional differences) and between the different regions (interregional differences) were analyzed. Regions are shown in hematoxylin-stained tissue sections in Supplementary Figure S11. Intraregional and interregional differences in expression level were analyzed by using a Mann–Whitney test and the Cohen effect size (65). For a carcinoma strongly expressing *HER2* (35 *HER2* copies, *HER2* IHC 3+), the interregional heterogeneity was weak with Cohen effect sizes ranging from $r = 0.09$ to $r = 0.11$ (Figure 4B, left, Supplementary Table S1). The intraregional differences within regions 2 and 3 were low, while region 1 showed a slightly increased heterogeneity in *HER2* expression levels (Figure 4B right). The carcinoma section with a lower *HER2* protein expression status (five *HER2* copies, *HER2* IHC 2+) showed weak interregional heterogeneity between regions 2 and 3 ($r = 0.04$) and moderate to strong differences in *HER2*

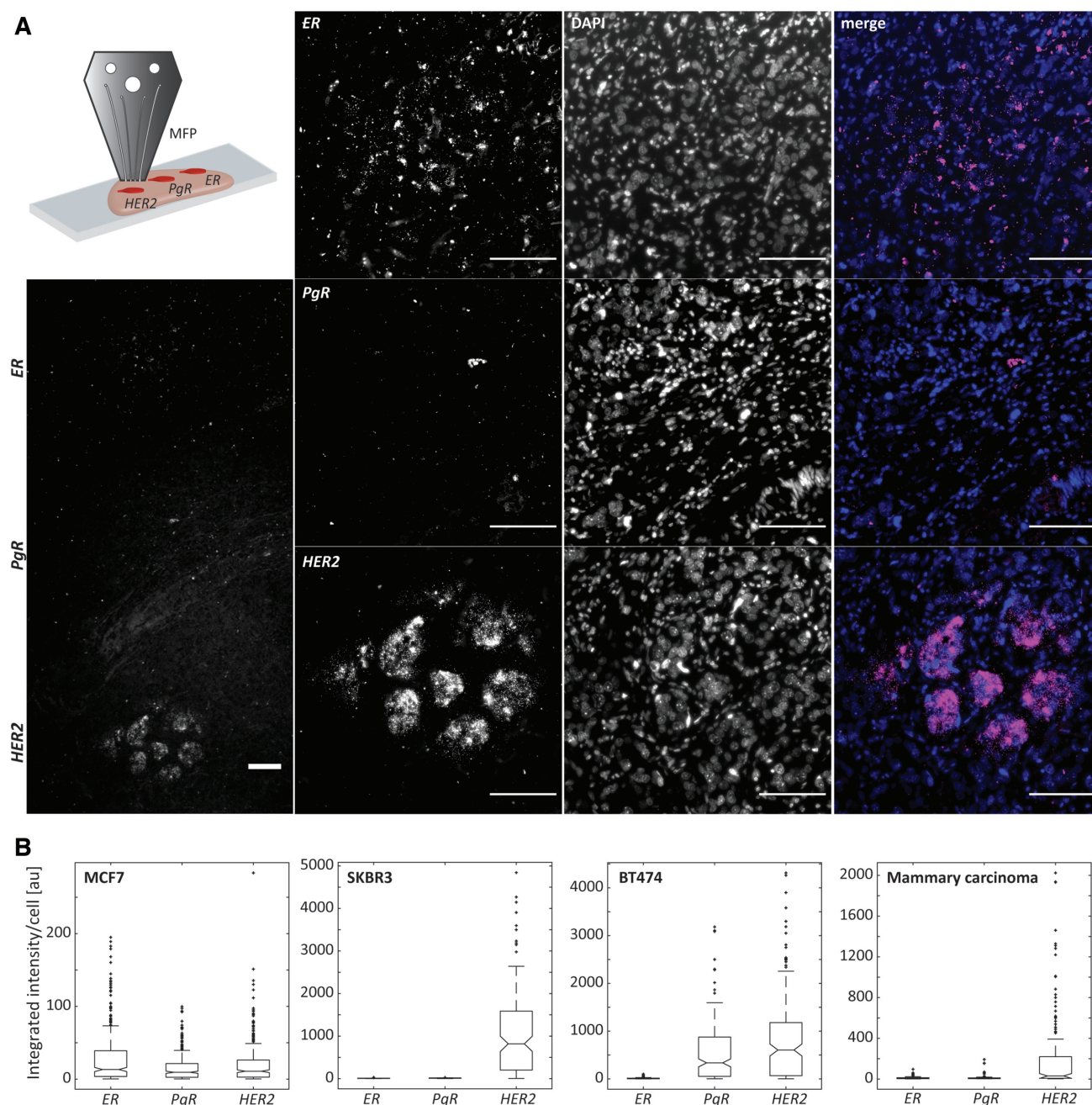


Figure 3. Single color multiplexed RNA-ISH for the detection of breast cancer biomarkers. (A) The primary probes for *ER*, *PgR*, and *HER2* were delivered to spatially distinct regions of a mammary carcinoma section using a microfluidic chip (schematic in upper left corner). Fluorescence images of the signal detected for the breast cancer biomarkers *ER*, *PgR* and *HER2* in a mammary carcinoma section. Scale bar: 100 μ m. (B) Fluorescence intensity of FISH signals detected for *ER*, *PgR* and *HER2* integrated per sectioned cell for FFPE sections of the breast cancer cell lines MCF7, SKBR3 and BT474 (see Supplementary Figure S7) as well as the mammary carcinoma from (A). 500 ms excitation.

expression levels when comparing them to region 1 ($r = 0.36$ and $r = 0.34$ for regions 1 and 2, and 1 and 3, respectively) (Figure 4C). Additionally, intraregional heterogeneity was moderate for region 1 with effect sizes ranging from $r = 0.05$ to $r = 0.4$ between different areas within the region. These results show that the molecular heterogeneity of the *HER2* status differs between tumors and patients. While some tumors display a homogeneous distribution of *HER2* expression levels, others show moderate to large variations. His-

tological and pathological evaluations of the tissue sections corresponded to the *HER2* expression patterns observed using RNA-ISH. The highly *HER2*-overexpressed tissue section (Figure 4B) contained 35 copies of the *HER2* gene and was classified as *HER2* 3+ when using immunohistochemistry, while the low *HER2* expressing carcinoma carried 5 *HER2* gene copies and had a *HER2* score of 2+. The evaluation of this tumor tissue was equivocal. Interestingly, these equivocal results corresponded to a tissue with moderate to

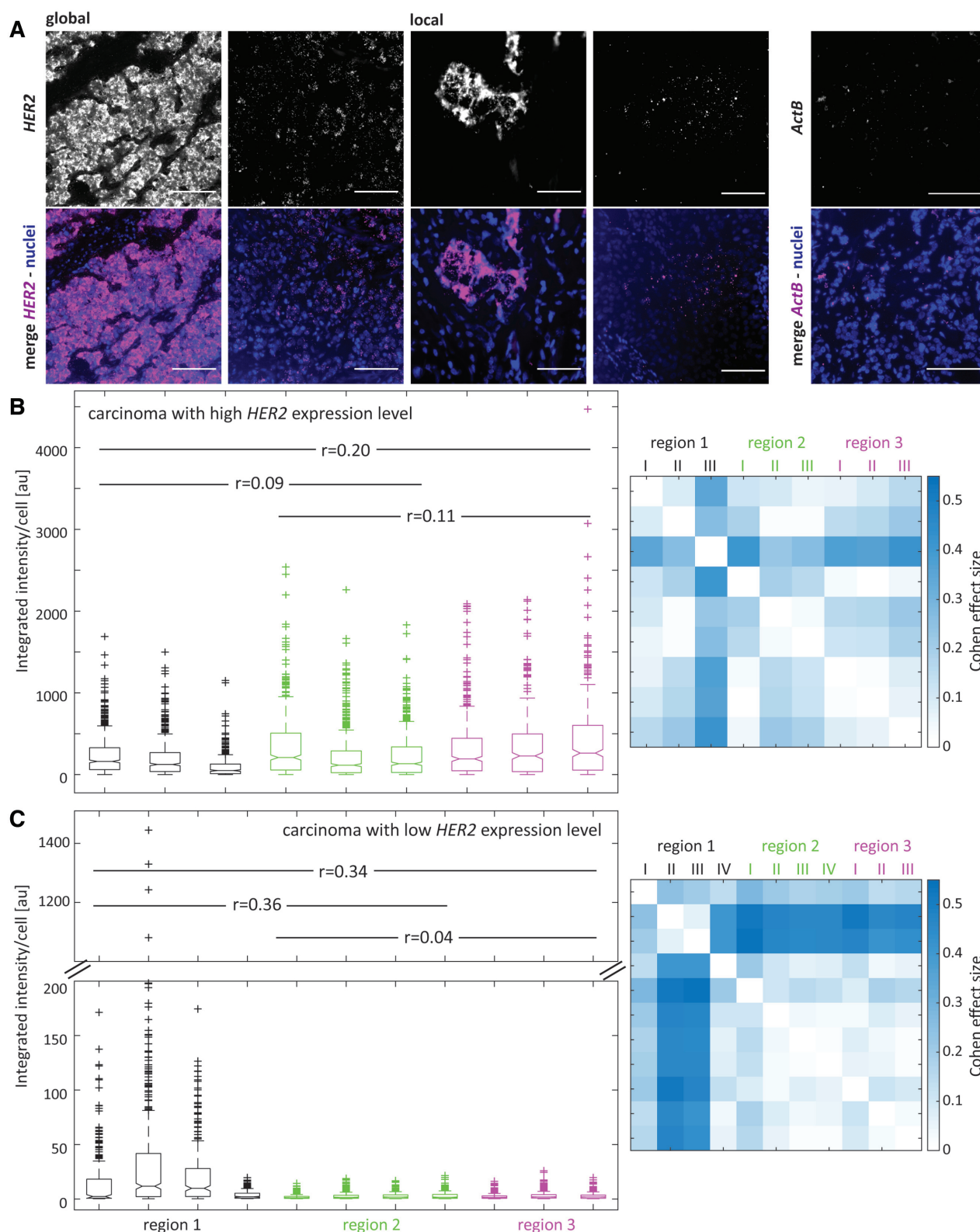


Figure 4. Spatially multiplexed RNA-ISH to detect spatial tumor heterogeneity in mammary carcinoma sections. (A) Fluorescence images of RNA-ISH signals detected for *HER2* and *ActB* by global ISH or local ISH experiments on mammary carcinoma sections. Nuclei visualized using DAPI. The images display the inter- and intratumor heterogeneity of the *HER2* expression. Scale bar: 100 μ m. (B, C) Integrated fluorescence intensity of the *HER2* signal per sectioned cell in three different regions of a strongly (B) (35 *HER2* copies, *HER2* IHC 3+) and low (C) (five *HER2* copies, *HER2* IHC 2+) *HER2*-expressing carcinoma tissue section (left). Using a Mann–Whitney test, the Cohen effect size (r) for interregional heterogeneity was calculated from the combined areas (I–IV) within a region and displayed by the values given in the graph on the left. A matrix of Cohen effect sizes calculated for the individual areas within or between regions (intra- and interregional heterogeneity) is shown on the right. Corresponding regions are shown on hematoxylin and immunohistochemically stained tissue sections in Supplementary Figure S11A and B.

high spatial heterogeneity in *HER2* transcript levels, suggesting the importance of molecular tumor heterogeneity for the differentiation of breast carcinomas into subtypes.

Multiplexed analysis revealed heterogeneity in *HER2* gene expression in relation to the expression levels of *ActB* (Figure 5A). While only a low variation in the expression of *ActB* was detected between three different regions of the tissue ($r = 0.08$ – 0.21), the expression level of *HER2* was significantly lower in region 3 compared to the other two regions ($r = 0.74$ for both, comparison between regions 1 and 3, and between regions 2 and 3). This suggests a differential regulation of *HER2*, while the overall RNA content between the regions is constant. A comparison of the *HER2/ActB* expression ratios showed an expression of *HER2* in region 3 that was 15 times lower than in both other regions analyzed (*HER2/ActB* (region 1) = 0.7, *HER2/ActB* (region 2) = 0.6, *HER2/ActB* (region 3) = 0.04).

Correlation between transcript and protein expression

We further investigated the correlation between differences in expression at the transcript and protein level. We determined *HER2* expression levels and their intraregional and interregional differences of a carcinoma tissue section with *HER2* status of 3+ in immunohistochemistry results (Figure 5B and C, Supplementary Table S1). *HER2* protein levels were analyzed using standard immunohistochemistry and quantitative immunohistochemistry (Figure 5B and D) (24). Between the four regions analyzed, the heterogeneity of *HER2* gene expression showed a small effect size between regions 1, 2, and 4. It was moderate when comparing these regions to region 3 ($r = 0.26$ for regions 1 and 3 as well as 4 and 3, and $r = 0.19$ for regions 2 and 3). Correspondingly, *HER2* protein showed a higher expression in region 3 when compared to the other regions (Figure 5C). Evidently, we observed a correlation in the spatial heterogeneity of *HER2* between the transcript and protein levels.

In summary, spatially multiplexed RNA-ISH can be used to detect spatial molecular tumor heterogeneity of a panel of biomarkers or in combination with house-keeping genes and internal controls. The transcript level can thereby provide additional quantitative information about the gene expression status and may be used in a complementary fashion to existing approaches for biomarker analysis, being especially advantageous for equivocal tumor cases.

DISCUSSION

Spatial tumor heterogeneity is an important driver in tumor progression and resistance development and is currently rarely considered in clinical decision making. Therefore, a feasible approach describing molecular tumor heterogeneity is needed. A variety of research methods to study spatial cellular heterogeneity at the protein and transcript expression level exists. Multiplexed immunohistochemistry, which is commercially available as Opal and UltiMapper detection schemes for automated platforms, allows to spatially detect up to 16 different proteins (14,70–72). Recent developments additionally explore the label-free detection of biomarkers using infrared imaging (73–76) or deep learning approaches on histological images (77–80). Furthermore,

a multitude of new techniques for the spatial analysis of transcripts have recently been developed. They range from location-inference for single-cell sequencing data (81–85) or next-generation sequencing of laser capture microdissected sections (47–49) to *in situ* sequencing (41–44) or patterned microarrays (45,46). Depending on the application, each of the methods has its own advantages, such as the number of different transcripts detectable, the efficiency of detection, or the quantification of gene expression. However, many of these methods require the use of live cells or fresh frozen tissue to ensure a stable quality of RNA. As FFPE samples are one of the most common sample types in pathology laboratories and tissue biobanks (86), a method to detect gene expression levels despite formalin-fixation is highly advantageous and may lead to translation of the technique to other laboratories. To limit the technical complexity and make spatial transcript analysis potentially accessible for clinical use, we therefore developed a novel method for multiplexed RNA-ISH, which has been shown to be compatible with FFPE tissues (50–52). RNA-ISH provides the additional advantage of detecting transcripts with low expression levels, such as non-coding RNAs. The method is based on existing RNA detection schemes that use amplification and brightfield or single-color fluorescence detection. We leveraged the microfluidic chip's ability to precisely and locally deliver chemicals (59,60) to spatially confine primary ISH-probes on FFPE sections and thus achieve spatial multiplexing with simple detection schemes.

By using spatial multiplexing, the number of different targets for amplification-based RNA-ISH techniques can be increased. Current non-fluorescent and enzyme-based amplification approaches for ISH have the advantage of abolishing the need for highly sensitive detection optics (56). However, they are limited to two different target samples. By applying spatial multiplexing, the number of different targets can easily be scaled and is only limited by the number of positions available in a homogeneous region of the tissue section. The established method provides the additional advantage that signal intensities within a section can be directly compared without the need to control for variations in pretreatment conditions or saturation of the amplification reaction. As multiplexing using a single color is possible, differences in quantum yield, excitation power, and excitation and emission filter bandwidth do not have to be corrected for. Furthermore, the spectral multiplexing of transcripts may be advantageous when assessing highly expressed genes, where the dense distribution hinders hybridization to multiple targets.

We demonstrated the utility of the technique by the detection of *HER2* transcripts with tissue internal positive and negative controls and the analysis of the predictive breast cancer biomarker panel *ER/PgR/HER2* in a single tumor tissue section. In clinical decision making, control reactions are extremely important as they provide information about tissue processing conditions, integrity of the molecular content, and performance of the analytical assay. Internal controls are the most precise measure and are therefore preferred over external controls on consecutive tissue sections or other samples (87). Including internal controls in RNA-ISH facilitates the evaluation of RNA integrity and stability of the experimental conditions and is thus

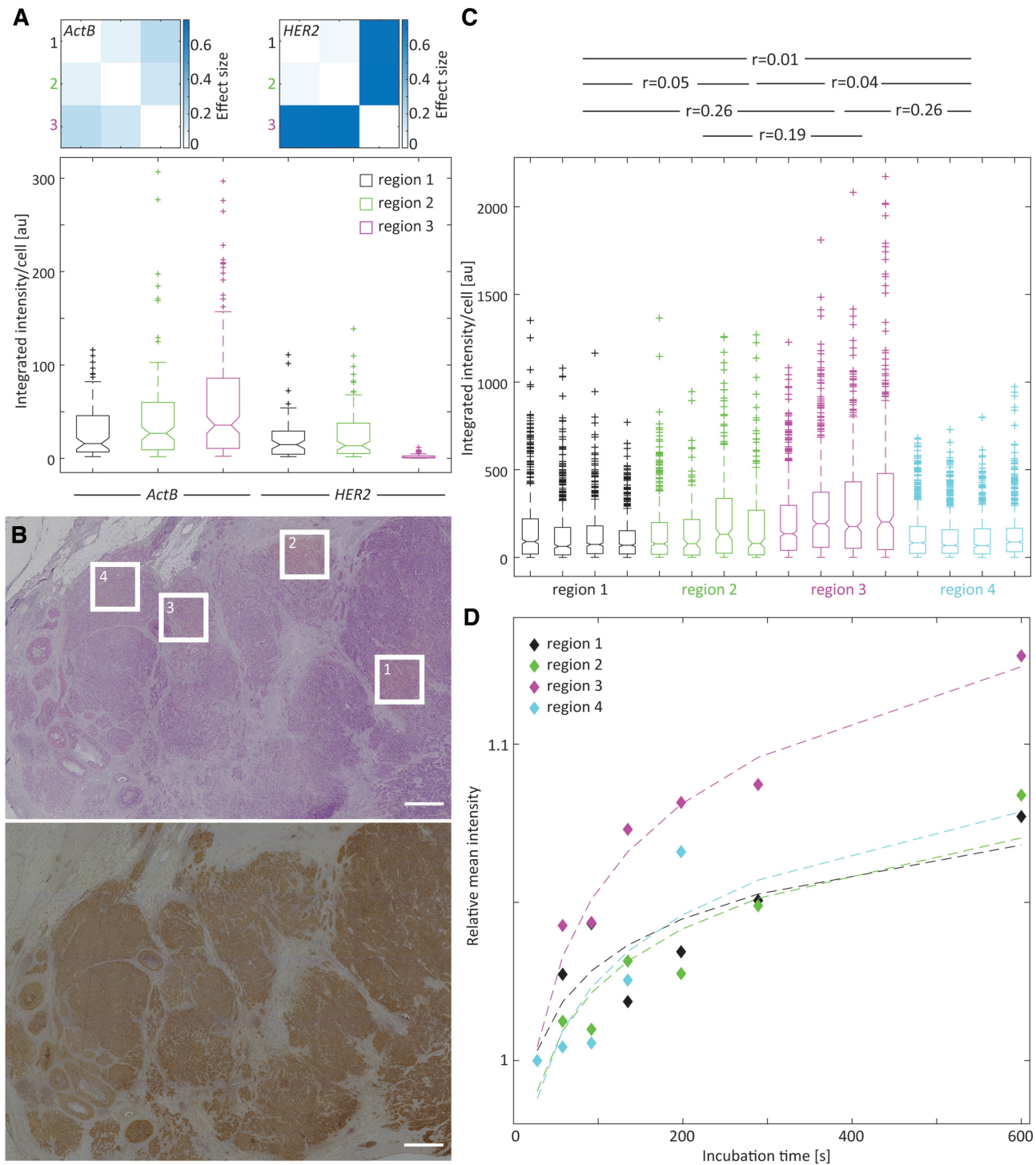


Figure 5. Differential *HER2* gene expression in mammary carcinoma sections. (A) Integrated RNA-ISH fluorescence intensity per cell section for *ActB* and *HER2* in a mammary carcinoma section (lower image). Cohen effect sizes describing interregional differences in expression levels calculated using a Mann-Whitney test are shown as matrices for *ActB* and *HER2*, respectively (upper images). (B) Histological overview of the tumor tissue section stained with hematoxylin and eosin (upper image) and *HER2* immunohistochemistry (lower image) with the regions 1–4 used for local RNA-ISH detection in (C). Scale bar: 1 mm. (C) Integrated RNA-ISH fluorescence intensity (*HER2*) per cell section for four areas in each of the four different tumor regions in a mammary carcinoma section with a *HER2* IHC status of 3+. The entire tissue sections including the regions of interest are shown as hematoxylin and IHC stained tissues in Supplementary Figure S11C. The Cohen effect size (r) for interregional heterogeneity was calculated from all areas (1–4) within each region combined by using a Mann-Whitney test and its values are displayed above the graph. (D) Quantitative immunohistochemistry using a gradient of primary antibody incubation times by local antibody delivery (24). The graph shows the relative mean intensity of the diaminobenzidine (DAB) signal after the amplification of the secondary antibody signal through a horseradish peroxidase reaction versus the incubation time of the primary antibody. It provides a quantitative measure of protein expression levels.

an important measure for assay performance. The established assay can thereby aid in implementing these internal controls.

The information from the multiplexed RNA-ISH analysis of panels of biomarkers can be used to allocate breast cancer cell pellet sections to their molecular subtype (MCF7: luminal A, BT474: luminal B, SKBR3: HER2-overexpressed (67)). Complementary to protein expression analysis, the same information derived from transcript expression levels can therefore be used to classify tumor tissue sections into their molecular subtypes. This is in accordance with the results by Alba *et al.* (2012), who have shown a 95.2% correlation of gene expression in RNA-ISH compared to immunohistochemistry and a 96.5% concordance of RNA-ISH with gene amplification by DNA-FISH (63). Baehner *et al.* (2010) additionally investigated the correlation between the RNA expression levels of *HER2* detected by qRT-PCR and DNA-FISH. The observed concordance of 97% between the two methods shows that RNA expression levels are a reliable indicator for patient stratification (88), a finding further confirmed by Vassilakopoulou *et al.* (89), Wang *et al.* (2), and Wu *et al.* (90). Although multiplexing with spectral unmixing and AQUA analysis enable a certain degree of quantitation of immunohistochemistry and immunofluorescence results (14,18,22,23), the quantification of immunohistochemistry results is still not precise and leads to 32% equivocal results, 84% of which can be resolved by subsequent DNA-FISH (2,63). The analysis of expression at the RNA level presents an alternative approach with the additional advantage of using oligonucleotide probes over antibodies, many of which are not suitable for obtaining reproducible IHC results on FFPE tissue (91,92). For PD-L1 testing, for example, different antibodies, assays, and evaluation criteria are used to predict PD-1/PD-L1 inhibitor therapy (93). Through using RNA-ISH, the standardization of PD-L1 testing may be facilitated. The clinical application of approaches such as Oncotype DX, 70-gene signature, or PAM-50 have illustrated the usefulness of analyzing transcript expression levels for breast cancer prognosis and prediction (26–29,94,95). Further studies showed the advantages of using mRNA expression profiles for the therapeutic stratification of patients with breast cancer (TAILORx (96), MINDACT (27), EndoPredict (97,98)). When using these techniques, the relative quantification of gene expression levels is achieved by using PCR and microarrays, while the spatial information is not preserved. Therefore, combinations of quantitative gene expression analysis at the transcript level with spatially resolved DNA-FISH or immunohistochemistry can reveal additional important information on molecular heterogeneity. Alternatively, RNA-ISH offers the possibility of the relative quantification of transcripts directly in a spatially defined manner on a single tissue section. Annaratone *et al.* quantified *ER* and *HER2* transcripts in FFPE tissue sections and successfully stratified the tissues into molecular subtypes (99). Using the method described in this paper, spatial differences of multiple transcripts of interest can be assessed on a single tissue section. We used the approach to evaluate the spatial heterogeneity of *HER2* transcripts and observed varying interregional and intraregional differences in the gene expression level for the carcinoma sec-

tions investigated. Interestingly, we observed heterogeneity in *HER2* transcript levels with moderate to strong effect size in carcinoma tissue with equivocal results from *HER2* gene copy number analysis and immunohistochemistry. This suggests that spatial heterogeneity contributes to the significance of molecular subtype differentiation, making it an even more important parameter to consider in clinical decisions. Especially in equivocal cases, spatially defined transcript levels can provide a complementary approach and additional indicator for predictive and prognostic decisions. We observed a correlation of interregional differences in the *HER2* expression between the transcript and protein level. This is in accordance with previous results revealing spatial heterogeneity of *HER2* breast cancer on the transcript or protein level (24,99,100).

We believe that the local delivery of probes and antibodies will facilitate the multimodal detection of RNA and proteins on single tissue sections by combining RNA-ISH and immunohistochemistry. Previous studies have shown the simultaneous detection of transcripts and proteins on a single monolayer of cells or tissue section (101–103) and recent elaborate techniques combine RNA-ISH and immunofluorescence with metal-based detection using mass cytometry (104). Furthermore, owing to its immense flexibility, the microfluidic chip may be used for the simultaneous local delivery of *in situ* detection moieties (e.g. hybridization probes, antibodies) and local removal of cells for genetic or transcriptomic analysis (61,105). We can thus imagine its capabilities to provide spatial information about whole genomes, transcriptomes, or even proteomes. The quantitation of transcripts and proteins in a spatially defined manner holds great promise for the elucidation of tumor development and progression. Future studies on cohorts of tumor biopsies can provide information on the applicability of the technique for clinical interpretation.

To summarize, our work provides a versatile approach for multiplexed *in situ* detection of transcripts using single-color or brightfield detection. As methods that recover spatial information provide the possibility to investigate dynamic processes, development of lineages, connectivity, and alterations in tissue architecture in development and disease, we believe that the developed technique will find application in a wide variety of biological and medical contexts.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

We thank Susanne Dettwiler for generating and providing tumor tissue sections. We are grateful to Daniel Schulz and Bernd Bodenmiller for their advice in the initial phase of our experiments and to Szymon Stoma and Simon F. Norrelykke for the discussion on image data analysis. Thanks are due to Robert D. Lovchik for his advice on imaging. We thank Natascha Brinskelle-Schmal for providing feedback on initial experiments. L.V. wishes to thank Alexander Borodavka and Waldemar Schrimpf for in-depth scientific discussions. We are grateful to Linda Rudin for valuable comments on the manuscript. We thank Emmanuel Dela-

marche, Walter Riess, and Heike E. Riel for their continuous support.

FUNDING

European Research Council [ERC 727761 (BioProbe-PIT) to G.V.K.]. Funding for open access charge: European Research Council.

Conflict of interest statement. L.V., A.F.K., D.H. and G.V.K. were employed by IBM Research GmbH and P.S. was employed by Universitätsspital Zürich while working on this project. The authors have no conflict of interest.

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